Changes in relaxin production by the theca during the preovulatory period of the pig

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Summary. Theca and granulosa layers were isolated from the preovulatory follicles of prepubertal gilts which were untreated (Group A), killed 72 h after 1000 i.u. PMSG (Group B), killed 84 h after PMSG (Group C), or killed 84 h after PMSG + 500 i.u. hCG given at 72 h (Group D). The tissues from individual follicles were cultured for 24 h alone (C), with FSH (F) or with LH (L), and the content of immunoreactive relaxin in the culture media was measured by RIA. Concentrations of relaxin-like material were close to the limit of detection of the assay in all granulosa cell cultures, and in the thecal cultures from the untreated gilts. However, thecal cultures from all 19 treated gilts produced relaxin. The mean ± s.e.m. concentrations (pg/follicle) in Groups AC, BC, CC and DC were 26·5 ± 3·04, 93·1 ± 4·6, 138 ± 16·4 and 285·6 ± 54·1 respectively. Therefore relaxin production was stimulated by PMSG \( (P < 0·05) \), with hCG treatment \textit{in vivo} leading to a further significant increase \( (P < 0·05) \). 

\textit{In-vitro} treatment with gonadotrophins had no effect in Groups A, C and D, but in Group-B gilts LH produced a significant \( (P < 0·05) \) rise in relaxin levels. These studies indicate that the theca is the principal source of relaxin in the porcine preovulatory follicle. The increased production before ovulation suggests that relaxin may be involved in follicular growth or rupture.

Introduction

Relaxin is present in the follicular fluid of pregnant and non-pregnant sows (Bryant-Greenwood, Jeffrey, Ralph & Seamark, 1980; Matsumoto & Chamley, 1980). Its presence there can be accounted for partly by diffusion from adjacent corpora lutea, the commonly recognized source of the hormone. However, the same two studies have shown that segments of follicle wall, but not isolated granulosa cells, release relaxin \textit{in vitro}, suggesting that follicular biosynthesis of relaxin can occur. Small amounts of relaxin are produced by porcine granulosa cells cultured for several days in the presence of LH (Loeken, Channing, D'Eletto & Weiss, 1983). Relaxin immunoreactivity has also been detected in human plasma around the time of ovulation in patients subjected to ovarian hyperstimulation with HMG (Thomas, Loumaye & Ferin, 1980).

The present study was conducted (1) to determine the source of the relaxin in porcine follicles and (2) to see whether relaxin production by the follicle changed during the preovulatory period as

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this might provide some indication of the role of relaxin in the non-pregnant pig. The prepubertal
gilt has proved useful for the study of steroid biosynthesis by the isolated theca and granulosa
(Evans, Dobias, King & Armstrong, 1981) and we therefore used a similar system to examine
relaxin production during follicular development and maturation.

**Materials and Methods**

**Animals**

Thirty-two (32) prepubertal Yorkshire crossbred gilts weighing 90.2 ± 2.3 kg (mean ± s.e.m.)
were allocated randomly to four treatment groups: Group A, control, slaughtered without
treatment; Group B, slaughtered 72 h after i.m. injection of 1000 i.u. PMSG (Equinex: Ayerst
Laboratories, Montreal); Group C, slaughtered 84 h after 1000 i.u. PMSG; Group D, slaughtered
at 84 h after injection of 1000 i.u. PMSG, with 500 i.u. hCG (A.P.L.: Ayerst) administered i.m. at
72 h. The treatment for Group D normally results in ovulation ~ 42 h after hCG, with a small
degree of superovulation (Baker & Coggins, 1968; Baker, Mellish & Segal, 1969), of meiotically
mature oocytes from steriodogenically active follicles (Ainsworth, Tsang, Downey, Marcus &

**Culture technique**

Ovaries were collected immediately after slaughter, washed and kept on ice in Eagle’s Minimal
Essential Medium buffered with Hepes. Theca tissue (interna + externa) and granulosa cells were
isolated and cultured for 24 h as described previously (Evans et al., 1981) with the following
differences. Tissue was cultured in 1 ml medium containing no hormones (control), 0.5 µg FSH
(NIH–FSH–S11) or 0.5 µg LH (NIH–LH–B4); doses of hormones in this range are known to
stimulate steroid production in vitro in these circumstances (Evans et al., 1981). Each in-vitro
treatment was replicated 4 times. One follicle-equivalent of tissue (theca or granulosa separately)
was added to each culture well, except for some animals from which < 16 follicles were obtained; in
such cases, the same degree of minor replication was maintained by adding proportionately less
tissue to each culture dish and correcting the results of relaxin analyses accordingly. Samples of
medium were cultured with hormones but without tissue, as appropriate controls. Samples of tissue
were reserved without culture for estimation of initial content of relaxin. These were acidified with
0.2 ml 50% acetic acid and freeze-dried. Dissections were complete and culture begun within
8–10 h of slaughter.

Immediately after culture, media were collected and kept at −20°C. All samples were freeze-
dried for transportation to England for assay.

**Relaxin radioimmunoassay**

Porcine relaxin was measured by an homologous radioimmunoassay as described by Taverne et
al. (1982). The highly-purified relaxin standard (CMy) used in this assay was extracted from the
ovaries of pregnant sows (Taverne et al., 1982). Freeze-dried samples of culture medium were
reconstituted in assay buffer (0.05 M-sodium barbitone pH 8.5 containing 50% inactivated horse
serum) and were measured in triplicate at two dilutions. These showed parallelism with the
standard curve (Text-fig. 1). For the acidified tissue samples, 0.8 ml assay buffer was added to the
freeze-dried material and vortexed and the supernatant was assayed as above. The sensitivity of the
assay was 4.5 pg per tube. The inter- and intra-assay coefficients of variation were 13.7% and 6.3%
respectively. Freeze drying of samples was previously shown not to alter their relaxin content.
**Text-fig. 1.** Porcine relaxin radioimmunoassay standard curve: — standard preparations, —— 3 different medium samples from thecal cultures (□, ○, △), each assayed at two dilutions.

**Protein determination**

Freeze-dried tissue samples were reconstituted in 1 ml 1% sodium dodecyl sulphate and the protein in each sample was measured by the method of Lowry, Rosebrough, Farr & Randall (1951) using bovine serum albumin as a standard.

**Statistical analyses**

Ovaries were rejected if they had not responded to PMSG treatment (infantile, or did not possess large follicles) or if they had signs of previous ovulations. In addition there was accidental loss of some samples. In combination, this resulted in data being obtained from 7 animals in Group A, 4 in Group B, 8 in Group C and 7 in Group D.

The data were analysed for theca and granulosa separately by analysis of variance, using an unbalanced design. A logarithmic transformation was required to eliminate heterogeneity of variance but, for convenient appraisal, the data are presented in untransformed form. Duncan’s Multiple Range Test was used to compare specific main effect means. A probability level of \( P < 0.05 \) was regarded as significant.
Results presented herein represent accumulation of relaxin in medium per follicle equivalent of tissue over a 24-h culture period. Samples of medium 'blanks' containing FSH or LH but no tissue and also cultured for 24 h had relaxin assay values close to the limits of detection and no corrections were made. The initial content of relaxin in representative samples of tissue before culture is shown in the figures but has not been subtracted from total accumulation in the medium. The final tissue content of relaxin (after culture) was not measured; the accumulation of relaxin in medium may therefore represent less than the total amount synthesized in culture. The average protein content of representative follicles at 0, 72 and 84 h after PMSG injection was: for granulosa 259, 472 and 427 µg respectively; for theca 357, 657 and 744 µg respectively. Since protein content was not measured for each cultured tissue the results have not been corrected for slight variations in quantity of tissue per culture well.

Granulosa cell relaxin production

The production of relaxin by granulosa cells at each stage of follicular development is represented in Text-fig. 2(a). The initial content of relaxin in tissue was below the limit of detection of the assay for many samples and, in general, the relaxin accumulation in media was close to the limits of detection. There was no significant effect of in-vivo treatment on relaxin accumulation nor was there any effect of in vitro hormone treatment.

Theca tissue relaxin production

As shown in Text-fig. 2(b) there was a significant effect \( P < 0.001 \) of in-vivo treatment (Group) on relaxin production; there was an increase in production in Groups B and C compared with Group A controls (72 and 84 h after PMSG; \( P < 0.05 \)), with even greater production in Group D \( P < 0.05 \) due to administration of hCG in vivo at 72 h after PMSG. There was no overall effect of adding FSH or LH to culture in vitro, but in Group B LH increased relaxin production \( P < 0.05 \).

Discussion

Relaxin is generally considered to be a hormone produced by the corpus luteum during pregnancy. However, it has been isolated from the follicular fluid of pregnant and non-pregnant sows, and it is also present in animals with polycystic ovaries (Bryant-Greenwood et al., 1980; Matsumoto & Chamley, 1980). Bryant-Greenwood et al. (1980) showed that although follicular relaxin levels tend to be higher in ovaries containing corpora lutea, suggesting diffusion from this source, segments of follicle wall will continue to release relaxin into the medium over a 4-day culture period indicating that the follicle is itself capable of relaxin biosynthesis. Although Loeken et al. (1983) have reported low-level relaxin secretion by granulosa cells isolated from large porcine follicles and cultured for several days in the presence of LH, they needed to concentrate their medium before the relaxin levels were measurable and neither Matsumoto & Chamley (1980) nor Bryant-Greenwood et al. (1980) could detect relaxin in cultures of isolated porcine granulosa cells. We were likewise unable to measure relaxin production by the granulosa much in excess of the limits of detection of the assay, and there was no stimulation by PMSG or hCG in vivo or FSH or LH in vitro at doses which normally evoke steroidogenic responses.

Our study indicates for the first time that it is the theca rather than the granulosa which is the major source of relaxin in porcine follicles. We also show that there is a significant increase in relaxin production by the theca as the time of ovulation approaches. This was initially stimulated in
Text-fig. 2. In-vitro production of relaxin by porcine granulosa cells (a) and theca tissue (b), measured as accumulation of relaxin per follicle equivalent of tissue in media after 24 h culture in the presence of no added hormone (C), 0.5 µg FSH/ml (F) or 0.5 µg LH/ml (L). The initial content of relaxin in the tissue before culture is also shown (I). Some of these values were below the limit of detection of the assay in which case the limit of detection was used in calculating the means. The in-vivo treatments were: Group A, controls (no in-vivo treatment); Group B, slaughter 72 h after PMSG; Group C, slaughter 84 h after PMSG; and Group D, slaughter 84 h after PMSG with hCG at 72 h (see ‘Materials and Methods’).

In-vivo with PMSG, but a further increase in output in vitro occurred from ovaries after hCG injection in vivo. In-vitro treatment with LH was only effective 72 h after PMSG, whereas FSH treatment in vitro did not alter relaxin production at any of the times studied. It therefore remains unclear whether the in-vivo injections of PMSG and hCG acted directly to enhance relaxin release, or whether some intermediate factors were involved. For example, Evans et al. (1981; 1983) have shown that a similar treatment regimen alters both the steroid and prostaglandin secretion patterns of the theca and granulosa and these changes could in turn influence relaxin production.

As the follicular capacity to release relaxin increases near the time of ovulation it seems likely that relaxin is involved in this process (see Bryant-Greenwood, 1982). One possible role for relaxin could be in the dissolution of the follicle wall and the release of the cumulus oophorus. The tunica albuginea and the theca externa of the pig contains an extensive network of collagen fibrils (Corner, 1919; Espey, 1967a). In the hours preceding ovulation this framework undergoes physiological deterioration resulting in a marked decrease in the tensile strength of the follicle wall.
Therefore theca-derived luteal cells, found by Bryant-Greenwood, allow relaxation to bring about a dose-related rise in the secretion of plasminogen activator by cultured rat granulosa cells indicates that relaxin may be one of the many factors involved in the control of these events (Too, Weiss & Bryant-Greenwood, 1982).

Smooth muscle is present in the hilar and medullary regions of the mammalian ovary and it has been reported to occur in the theca externa of several species, including the sow (Corner, 1919; Guttmacher & Guttmacher, 1921). In their detailed study of the hamster ovary, Martin & Talbot, 1981a, b) show that the theca externa only contracts in the last few minutes before ovulation, when the base of the follicle is pushed upwards. Smooth muscle tends to contract in response to stretch, so that, by relaxing muscle cells in the theca externa during the preovulatory period, relaxin could allow the follicle to expand rapidly at this time whilst preventing premature contraction.

The classic work of Corner (1919) showed clearly that both the theca and granulosa contribute to the formation of the corpus luteum in the sow and this has been confirmed ultrastructurally (Cavazos et al., 1969). Both papers suggest that the granulosa layer gives rise to the large (granulosa) lutein cells, whereas the theca interna forms the small (theca) lutein cells. Three separate reports on the localization of relaxin within the corpus luteum of the pregnant sow all state that the peptide is found in granulosa lutein cells, whilst omitting any reference to the presence of theca lutein cells (Belt, Cavazos, Anderson & Melampy, 1971; Larkin, Fields & Oliver, 1977; Kendall, Plopper & Bryant-Greenwood, 1978). However, our results with cultured follicular tissue indicate that in the preovulatory follicle it is the theca rather than the granulosa which secretes relaxin. If the same cell line continues to make the same peptide hormone after ovulation this would imply that the small theca-derived cells can become large luteal cells. In 1966, following a study of the bovine corpus luteum Donaldson & Hansel wrote "The granulosa-derived luteal cells develop into functioning luteal cells by about Day 4 of the estrous cycle, after which they undergo no further divisions. Therefore granulosa-derived cells have limited potential. However the smaller luteal cells derived from the theca interna respond to plasma luteotropin, multiply and grow into large luteal cells. These are mainly responsible for the growth of the corpus luteum following Day 4". As no-one has yet labelled preovulatory theca or granulosa cells and followed their subsequent fate this hypothesis has not been tested. However, our data on relaxin suggest that it may be correct.

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References


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